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FILE 'HOME' ENTERED AT 19:57:49 ON 09 OCT 2002

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=> file caplus

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SINCE FILE

TOTAL

FULL ESTIMATED COST

ENTRY 0.21

SESSION 0.21

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FILE COVERS 1907 - 9 Oct 2002 VOL 137 ISS 15 FILE LAST UPDATED: 8 Oct 2002 (20021008/ED)

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CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

=> s oxid?

L1 2369625 OXID?

=> s l1 and aqueous

148293 AQUEOUS

1 AQUEOUSES

148294 AQUEOUS

(AQUEOUS OR AQUEOUSES)

933909 AQ

119 AQS

933982 AQ

(AQ OR AQS)

968682 AQUEOUS

(AQUEOUS OR AQ)

L2 169395 L1 AND AQUEOUS

=> s 12 catalyst

MISSING OPERATOR L2 CATALYST

The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

```
=> s 12 and catalyst
        601917 CATALYST
        585163 CATALYSTS
        767433 CATALYST
                  (CATALYST OR CATALYSTS)
         24536 L2 AND CATALYST
L3
=> s 13 and nanoparticle
         11088 NANOPARTICLE
         17675 NANOPARTICLES
         18714 NANOPARTICLE
                  (NANOPARTICLE OR NANOPARTICLES)
            67 L3 AND NANOPARTICLE
L4
=> s l4 and stabiliz?
        354798 STABILIZ?
             8 L4 AND STABILIZ?
L5
=> s 15 and alcohol
        165042 ALCOHOL
        124445 ALCOHOLS
        270252 ALCOHOL
                  (ALCOHOL OR ALCOHOLS)
        503294 ALC
        162712 ALCS
        587755 ALC
                  (ALC OR ALCS)
        681787 ALCOHOL
                  (ALCOHOL OR ALC)
             3 L5 AND ALCOHOL
L6
=> dis 16 1-3 bib abs
1.6
     ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS
     2002:90690 CAPLUS
ΑN
     136:285000
DN
     Deposition of Platinum Nanoparticles, Synthesized in
TΙ
     Water-in-Oil Microemulsions, on Alumina Supports
     Ingelsten, Hanna Haerelind; Beziat, Jean-Christophe; Bergkvist, Kristina;
ΑU
     Palmqvist, Anders; Skoglundh, Magnus; Hu, Qiuhong; Falk, Lena K. L.;
     Holmberg, Krister
     Competence Centre for Catalysis, Department of Applied Surface Chemistry,
CS
     Department of Applied Physics, and Department of Experimental Physics,
     Chalmers University of Technology, Goeteborg, SE-412 96, Swed.
SO
     Langmuir (2002), 18(5), 1811-1818
     CODEN: LANGD5; ISSN: 0743-7463
     American Chemical Society
PR
     Journal
DΤ
     English
LA
     Pt nanoparticles were prepd. in H2O-in-oil microemulsions and
AB
     deposited on .gamma.-alumina using 2 different methods. In the 1st
     method, the alumina support was added to the particle suspension and the
     microemulsion was subsequently destabilized by addn. of THe, whereby the
     particles were deposited on the alumina support. In the other method, the
     Pt nanoparticles were transferred to an aq. soln.
     where they were redispersed by a stabilizing surfactant prior to addn. of the alumina support. The size of the microemulsion droplets and
     of the unsupported Pt particles was in the range of a few nanometers as
     measured by a dynamic light scattering technique (photon correlation
     spectroscopy). The size of the unsupported Pt nanoparticles and
     of the particles deposited on alumina was studied by TEM. Both methods
     for Pt particle deposition resulted in some degree of particle
     agglomeration, the 1st probably because of too-fast destabilization of the
```

microemulsion and the 2nd due to inefficient redispersion of the Pt particles when transferred to the aq. phase. All samples studied showed high catalytic activity for CO oxidn. by oxygen. The highest activity was found for those samples prepd. via the redispersion method where a relatively weak interaction was achieved between the redispersed Pt particles and the alumina. RE.CNT 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS 2000:666737 CAPLUS 133:254142 Catalytic method for modifying carbohydrates, alcohols, aldehydes or polyhydroxy compounds Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter; Haji Begli, Alireza Sudzucker Aktiengesellschaft, Germany in tout PCT Int. Appl., 45 pp. CODEN: PIXXD2 Patent German FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE 20000921 WO 2000-EP2351 WO 2000055165 A1 20000316 W: AU, CA, IL, US RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE DE 19911504 A1 20001019 DE 1999-19911504 19990316 EP 2000-925117 EP 1165580 20020102 20000316 A1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI AU 747812 В2 20020523 AU 2000-43953 20000316 PRAI DE 1999-19911504 A 19990316 WO 2000-EP2351 W 20000316 Industrial conversion of the title compds. in aq. phase is carried out in the presence of metal catalysts consisting of polymer-stabilized nanoparticles. A catalyst of this type is not deactivated by the conversion reaction as long as the stabilizing interaction between the polymer and the nanoparticles is maintained. For example, activity of an Al203-supported, poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn. given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs. RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS 2000:566746 CAPLUS 134:117443 Hydrogenation of olefins in aqueous phase, catalyzed by ligand/protected and polymer-protected rhodium colloids Borsla, A.; Wilhelm, A. M.; Canselier, J. P.; Delmas, H. Laboratoire de Genie Chimique-UMR CNRS 5503 (INPT/UPS) Ecole Nationale Superieure d'Ingenieurs de Genie Chimique, Toulouse, 31078/4, Fr. Studies in Surface Science and Catalysis (2000), 130C(International Congress on Catalysis, 2000 Pt. C) 2002-2000 Congress on Catalysis, 2000, Pt. C), 2093-2098-CODEN: SSCTDM; ISSN: 0167-2991

PB Elsevier Science B.V.

DT Journal

L6

ΑN

DN

ΤI

ΙN

PΑ

SO

DT

LA

PΤ

AB

L6

ΑN DN

TI

ΑIJ CS

SO

LA English

AΒ M-tri-sulfonated triphenylphosphine oxide as its sodium salt

(OTPPTS) and highly water sol. polymers such as poly(vinyl alc.) (PVA) and poly(vinylpyrrolidone) (PVP) were used to stabilize colloidal suspensions of active rhodium particles. The stabilized colloids were used as catalysts in hydrogenation of oct-1-ene in a two-liq. phase system. The effect of various parameters on the stability and activity of the metal nanoparticles under more or less severe conditions was studied. For OTPPTS-protected rhodium colloid systems, the colloid stability improved by increasing the P/Rh molar ratio. Lowering the pressure also improved stability, but led to lower catalytic activity while temp. had no effect on stability but rather on the formation of the active species. The PVP(K15)-Rh colloids can be re-used more than one time at 50.degree. and 0.3 MPa without loss of activity, but not the PVP(K30)-Rh and PVA-Rh colloids. Recycling of the catalytic phase provided unchanged turnover frequencies. Suppression of co-solvent did not result in noticeable changes in activity, indicating that the reaction takes place at the interface. The MET micrographs of the catalytic phase of the OTPPTS-Rh system after use in hydrogenation shows 4 nm particles contg. rhodium oxide and metallic rhodium. IR spectroscopy anal. of the PVP-Rh colloids shows geminal and terminal Rh-CO species with a preponderance of the geminal ones.

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 15 and carbohydrate 108745 CARBOHYDRATE

> 116172 CARBOHYDRATES 176447 CARBOHYDRATE

> > (CARBOHYDRATE OR CARBOHYDRATES)

L7 1 L5 AND CARBOHYDRATE

=> dis 17 ibib abs

ANSWER 1 OF 1 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:666737 CAPLUS

DOCUMENT NUMBER: 133:254142

TITLE: Catalytic method for modifying carbohydrates

, alcohols, aldehydes or polyhydroxy compounds INVENTOR(S): Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf;

Vorlop, Klaus-Dieter; Haji Begli, Alireza

PATENT ASSIGNEE(S): Sudzucker Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 45 pp.

CODEN: PIXXD2

mutant DOCUMENT TYPE: Patent

German FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

LANGUAGE:

PATENT NO. KIND DATE APPLICATION NO. DATE -----WO 2000055165 A1 20000921 WO 2000-EP2351 20000316 W: AU, CA, IL, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE

DE 19911504 20001019 DE 1999-19911504 19990316 A1 EP 1165580 20020102 EP 2000-925117 A120000316

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI

AU 747812 B2 20020523 AU 2000-43953 20000316 PRIORITY APPLN. INFO.: DE 1999-19911504 A 19990316 W 20000316 WO 2000-EP2351

Industrial conversion of the title compds. in aq. phase is AB carried out in the presence of metal catalysts consisting of polymer-stabilized nanoparticles. A catalyst

of this type is not deactivated by the conversion reaction as long as the stabilizing interaction between the polymer and the nanoparticles is maintained. For example, activity of an Al2O3-supported, poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn. given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs.

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> dis hist

(FILE 'HOME' ENTERED AT 19:57:49 ON 09 OCT 2002)

FILE 'CAPLUS' ENTERED AT 19:58:09 ON 09 OCT 2002
L1 2369625 S OXID?
L2 169395 S L1 AND AQUEOUS
L3 24536 S L2 AND CATALYST
L4 67 S L3 AND NANOPARTICLE
L5 8 S L4 AND STABILIZ?
L6 3 S L5 AND ALCOHOL

L7 1 S L5 AND CARBOHYDRATE

=> s 15 and platinum

153377 PLATINUM

48 PLATINUMS

153387 PLATINUM

(PLATINUM OR PLATINUMS)

L8 2 L5 AND PLATINUM

=> dis 18 1-2 bib abs

L8 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2002 ACS

AN 2002:90690 CAPLUS

DN 136:285000

TI Deposition of **Platinum Nanoparticles**, Synthesized in Water-in-Oil Microemulsions, on Alumina Supports

- AU Ingelsten, Hanna Haerelind; Beziat, Jean-Christophe; Bergkvist, Kristina; Palmqvist, Anders; Skoglundh, Magnus; Hu, Qiuhong; Falk, Lena K. L.; Holmberg, Krister
- CS Competence Centre for Catalysis, Department of Applied Surface Chemistry, Department of Applied Physics, and Department of Experimental Physics, Chalmers University of Technology, Goeteborg, SE-412 96, Swed.

SO Langmuir (2002), 18(5), 1811-1818 deale | CODEN: LANGD5; ISSN: 0743-7463

- PB American Chemical Society
- DT Journal
- LA English
- AB Pt nanoparticles were prepd. in H2O-in-oil microemulsions and deposited on .gamma.-alumina using 2 different methods. In the 1st method, the alumina support was added to the particle suspension and the microemulsion was subsequently destabilized by addn. of THe, whereby the particles were deposited on the alumina support. In the other method, the Pt nanoparticles were transferred to an aq. soln. where they were redispersed by a stabilizing surfactant prior to addn. of the alumina support. The size of the microemulsion droplets and of the unsupported Pt particles was in the range of a few nanometers as measured by a dynamic light scattering technique (photon correlation spectroscopy). The size of the unsupported Pt nanoparticles and of the particles deposited on alumina was studied by TEM. Both methods for Pt particle deposition resulted in some degree of particle agglomeration, the 1st probably because of too-fast destabilization of the

microemulsion and the 2nd due to inefficient redispersion of the Pt particles when transferred to the aq. phase. All samples studied showed high catalytic activity for CO oxidn. by oxygen. The highest activity was found for those samples prepd. via the redispersion method where a relatively weak interaction was achieved between the redispersed Pt particles and the alumina. RE.CNT 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 2 OF 2 CAPLUS COPYRIGHT 2002 ACS 2000:666737 CAPLUS 133:254142 Catalytic method for modifying carbohydrates, alcohols, aldehydes or polyhydroxy compounds Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter;

Haji Begli, Alireza instant Sudzucker Aktiengesellschaft, Germany PΑ

SO PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DT Patent

LΑ German

FAN.CNT 1

L8

ΑN

DN

ΤI

IN

PATENT NO. KIND DATE APPLICATION NO. DATE PΙ WO 2000055165 A1 20000921 WO 2000-EP2351 20000316 W: AU, CA, IL, US RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE 20001019 DE 1999-19911504 19990316 DE 19911504 Α1 EP 1165580 Α1 20020102 EP 2000-925117 20000316 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI AU 747812 AU 2000-43953 20000316 В2 20020523

PRAI DE 1999-19911504 A 19990316 WO 2000-EP2351 W 20000316

AB Industrial conversion of the title compds. in aq. phase is carried out in the presence of metal catalysts consisting of polymer-stabilized nanoparticles. A catalyst of this type is not deactivated by the conversion reaction as long as the stabilizing interaction between the polymer and the nanoparticles is maintained. For example, activity of an Al203-supported, poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn. given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs.

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s hydrogenation 150203 HYDROGENATION 1908 HYDROGENATIONS L9 150451 HYDROGENATION (HYDROGENATION OR HYDROGENATIONS) => s 19 and aqueous 148293 AQUEOUS 1 AQUEOUSES 148294 AQUEOUS

(AQUEOUS OR AQUEOUSES)

933909 AQ 119 AQS 933982 AQ

(AQ OR AQS)

968682 AQUEOUS

(AQUEOUS OR AQ)

L10 14512 L9 AND AQUEOUS

=> s 110 and catalyst

601917 CATALYST

585163 CATALYSTS

767433 CATALYST

(CATALYST OR CATALYSTS)

L11 6442 L10 AND CATALYST

=> s 111 and metal

1371087 METAL

685692 METALS

1661953 METAL

(METAL OR METALS)

L12 1568 L11 AND METAL

=> s 112 and polymer (w) stabiliz?

849461 POLYMER

708926 POLYMERS

1158743 POLYMER

(POLYMER OR POLYMERS)

354798 STABILIZ?

3772 POLYMER (W) STABILIZ?

L13 3 L12 AND POLYMER (W) STABILIZ?

=> s 112 and polymer

849461 POLYMER

708926 POLYMERS

1158743 POLYMER

(POLYMER OR POLYMERS)

L14 89 L12 AND POLYMER

=> s 114 and nanoparticle

11088 NANOPARTICLE

17675 NANOPARTICLES

18714 NANOPARTICLE

(NANOPARTICLE OR NANOPARTICLES)

L15 5 L14 AND NANOPARTICLE

=> s 115 and aldehyde

89500 ALDEHYDE

84224 ALDEHYDES

139257 ALDEHYDE

(ALDEHYDE OR ALDEHYDES)

L16 0 L15 AND ALDEHYDE

=> s 115 and alcohol

165042 ALCOHOL

124445 ALCOHOLS

270252 ALCOHOL

(ALCOHOL OR ALCOHOLS)

503294 ALC

162712 ALCS

587755 ALC

(ALC OR ALCS)

681787 ALCOHOL

(ALCOHOL OR ALC)

L17 2 L15 AND ALCOHOL

=> dis 117 1-2 bib abs

- L17 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2002 ACS
- AN 2001:65241 CAPLUS
- DN 134:268058
- TI Synthesis and functionalities of poly(N-vinylalkyloamide). XIII. Temperature and pH dependence of the catalytic activity of colloidal platinum nanoparticles stabilized by poly[(vinylamine)-co-(N-vinylisobutyramide)]
- AU Chen, Chun-Wei; Arai, Kumiko; Yamamoto, Kazuya; Serizawa, Takeshi; Akashi, Mitsuru
- CS Dep. Applied Chem. Chemical Eng., Kagoshima Univ., Kagoshima, 890-0065, Japan
- SO Macromolecular Chemistry and Physics (2000), 201(18), 2811-2819 CODEN: MCHPES; ISSN: 1022-1352
- PB Wiley-VCH Verlag GmbH
- DT Journal
- LA English
- Colloidal platinum nanoparticles in the size range of 5-35 .ANG. AB have been successfully prepd. in water at room temp. by NaBH4 redn. of ionic platinum in the presence of poly[(vinylamine)-co-(Nvinylisobutyramide)] (PVAm-co-PNVIBA). The temp.- and pH-responsive copolymer was used for the first time as the stabilizer of colloidal metal particles. Three PVAm-co-PNVIBA copolymers with PVAm contents of 4.1, 8.3, and 19.8 mol-% were examd. The particle size and morphol. of the platinum colloids varied with the copolymer compn., as confirmed by TEM measurements. The polymer-stabilized Pt nanoparticles pptd. on heating above their crit. flocculation temps. (CFTs), which were strongly dependent on the soln. pH and the copolymer compn. The CFTs were 0.2-1.6.degree.C lower than the lower crit. soln. temps. (LCSTs) of the copolymers free in water and the differences increased with increasing PVAm content. The catalytic activity of the Pt nanoparticles was investigated in the aq. hydrogenation of allyl alc. It was found that the activity was regulated through temp. - and pH-induced phase sepn. The PVAm content also strongly effected the catalytic activity and the morphol. of phase sepd. catalysts. With a PVAm content of 4.1 mole, the colloidal platinum sol reversibly changed its catalytic activity with changes in temp.
- RE.CNT 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L17 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2002 ACS
- AN 2000:566746 CAPLUS
- DN 134:117443
- TI Hydrogenation of olefins in aqueous phase, catalyzed by ligand/protected and polymer-protected rhodium colloids
- AU Borsla, A.; Wilhelm, A. M.; Canselier, J. P.; Delmas, H.
- CS Laboratoire de Genie Chimique-UMR CNRS 5503 (INPT/UPS) Ecole Nationale Superieure d'Ingenieurs de Genie Chimique, Toulouse, 31078/4, Fr.
- SO Studies in Surface Science and Catalysis (2000), 130C(International Congress on Catalysis, 2000, Pt. C), 2093-2098
 CODEN: SSCTDM; ISSN: 0167-2991
- PB Elsevier Science B.V.
- DT Journal
- LA English
- AB M-tri-sulfonated triphenylphosphine oxide as its sodium salt (OTPPTS) and highly water sol. polymers such as poly(vinyl alc.)

 (PVA) and poly(vinylpyrrolidone) (PVP) were used to stabilize colloidal suspensions of active rhodium particles. The stabilized colloids were used as catalysts in hydrogenation of oct-1-ene in a two-liq. phase system. The effect of various parameters on the stability and activity of the metal nanoparticles under more or less severe conditions was studied. For OTPPTS-protected rhodium colloid systems, the colloid stability improved by increasing the P/Rh molar ratio. Lowering the pressure also improved stability, but led to lower

catalytic activity while temp. had no effect on stability but rather on the formation of the active species. The PVP(K15)-Rh colloids can be re-used more than one time at 50.degree. and 0.3 MPa without loss of activity, but not the PVP(K30)-Rh and PVA-Rh colloids. Recycling of the catalytic phase provided unchanged turnover frequencies. Suppression of co-solvent did not result in noticeable changes in activity, indicating that the reaction takes place at the interface. The MET micrographs of the catalytic phase of the OTPPTS-Rh system after use in hydrogenation shows 4 nm particles contg. rhodium oxide and metallic rhodium. IR spectroscopy anal. of the PVP-Rh colloids shows geminal and terminal Rh-CO species with a preponderance of the geminal ones.

RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 115 and carbohydrate 108745 CARBOHYDRATE 116172 CARBOHYDRATES 176447 CARBOHYDRATE (CARBOHYDRATE OR CARBOHYDRATES) 0 L15 AND CARBOHYDRATE L18 => s 115 and fructose 53219 FRUCTOSE 82 FRUCTOSES 53232 FRUCTOSE (FRUCTOSE OR FRUCTOSES) L19 0 L15 AND FRUCTOSE => s reduction 262772 REDUCTION 5081 REDUCTIONS 265419 REDUCTION (REDUCTION OR REDUCTIONS) 742417 REDN 39209 REDNS 766916 REDN (REDN OR REDNS) 891539 REDUCTION L20 (REDUCTION OR REDN) => s 120 and catalyst 601917 CATALYST 585163 CATALYSTS 767433 CATALYST (CATALYST OR CATALYSTS) 68034 L20 AND CATALYST L21 => s 121 and metal 1371087 METAL 685692 METALS 1661953 METAL (METAL OR METALS) L22 15952 L21 AND METAL => s 122 and nanoparticle 11088 NANOPARTICLE 17675 NANOPARTICLES 18714 NANOPARTICLE (NANOPARTICLE OR NANOPARTICLES) L23 129 L22 AND NANOPARTICLE

849461 POLYMER 708926 POLYMERS 1158743 POLYMER (POLYMER OR POLYMERS) L24 36 L23 AND POLYMER => s 124 and aldehyde 89500 ALDEHYDE 84224 ALDEHYDES 139257 ALDEHYDE (ALDEHYDE OR ALDEHYDES) L25 O L24 AND ALDEHYDE => s 124 and carbohydrate 108745 CARBOHYDRATE 116172 CARBOHYDRATES 176447 CARBOHYDRATE (CARBOHYDRATE OR CARBOHYDRATES) 0 L24 AND CARBOHYDRATE L26 => s 124 and alcohol 165042 ALCOHOL 124445 ALCOHOLS 270252 ALCOHOL (ALCOHOL OR ALCOHOLS) 503294 ALC 162712 ALCS 587755 ALC (ALC OR ALCS) 681787 ALCOHOL (ALCOHOL OR ALC) L27 6 L24 AND ALCOHOL => dis 127 1-6 bib abs ANSWER 1 OF 6 CAPLUS COPYRIGHT 2002 ACS 1.27 2001:65241 CAPLUS AΝ DN 134:268058 Synthesis and functionalities of poly(N-vinylalkyloamide). XIII. ТT Temperature and pH dependence of the catalytic activity of colloidal platinum nanoparticles stabilized by poly[(vinylamine)-co-(Nvinylisobutyramide)] ΑU Chen, Chun-Wei; Arai, Kumiko; Yamamoto, Kazuya; Serizawa, Takeshi; Akashi, Mitsuru CS Dep. Applied Chem. Chemical Eng., Kagoshima Univ., Kagoshima, 890-0065, Macromolecular Chemistry and Physics (2000), 201(18), 2811-2819 SO CODEN: MCHPES; ISSN: 1022-1352 PB Wiley-VCH Verlag GmbH DTJournal LA English AB Colloidal platinum nanoparticles in the size range of 5-35 .ANG. have been successfully prepd. in water at room temp. by NaBH4 redn . of ionic platinum in the presence of poly[(vinylamine)-co-(Nvinylisobutyramide)] (PVAm-co-PNVIBA). The temp.- and pH-responsive copolymer was used for the first time as the stabilizer of colloidal metal particles. Three PVAm-co-PNVIBA copolymers with PVAm contents of 4.1, 8.3, and 19.8 mol-% were examd. The particle size and morphol. of the platinum colloids varied with the copolymer compn., as confirmed by TEM measurements. The polymer-stabilized Pt nanoparticles pptd. on heating above their crit. flocculation temps. (CFTs), which were strongly dependent on the soln. pH and the copolymer compn. The CFTs were 0.2-1.6.degree.C lower than the lower crit. soln. temps. (LCSTs) of the copolymers free in water and the

differences increased with increasing PVAm content. The catalytic activity of the Pt nanoparticles was investigated in the aq. hydrogenation of allyl alc. It was found that the activity was regulated through temp.— and pH-induced phase sepn. The PVAm content also strongly effected the catalytic activity and the morphol. of phase sepd. catalysts. With a PVAm content of 4.1 mol%, the colloidal platinum sol reversibly changed its catalytic activity with changes in temp.

RE.CNT 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L27 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2002 ACS
- AN 2000:492945 CAPLUS
- DN 133:95064
- TI Preparation and catalysis of **polymer**-protected coinage **metal** nanoclusters
- AU Shiraishi, Yukihide; Hirakawa, Kazutaka; Toshima, Naoki
- CS Department of Materials Science and Engineering, Science University of Tokyo in Yamaguchi, Onoda-shi, Yamaguchi, 756-0884, Japan
- SO Kobunshi Ronbunshu (2000), 57(6), 346-355 CODEN: KBRBA3; ISSN: 0386-2186
- PB Kobunshi Gakkai
- DT Journal
- LA Japanese
- Prepn., characterization and catalysis of polymer-protected AB coinage metal nanoclusters were investigated, esp. from the viewpoint of bimetallization. Poly (N-vinyl-2-pyrrolidone) (PVP)-protected Au/Rh bimetallic nanoclusters with an av. diam. of 2.6 nm were prepd. by alc. redn. of HAuCl4 and RhChI in the presence of PVP. The Au/Rh bimetallic nanoclusters having an Au-core/Rh-shell structure work as more active ${\bf catalysts}$ for visible-light-induced hydrogen generation from water than the corresponding Au and Rh monometallic nanoclusters. Ag nanoclusters protected by poly(sodium acrylate) (PSA) were prepd. by UV irradn. of an alc.-water soln. of silver perchlorate in the presence of PSA and were applied to the **catalyst** for oxidn. of ethylene. The activity of Ag nanoclusters thus prepd. remarkably increases with increasing the reaction temp. Addn. of cesium and rhenium ions increases the catalytic activity of PSA-Ag nanoclusters as well. Thus, PSA plays an important role in this system. PVP-protected Cu/Pd alloy nanoclusters were prepd. by redn. in glycol at 198.degree.C, revealing high activity as a catalyst for hydration of acrylonitrile to acrylamide as well as for partial hydrogenation of 1,3-cyclooctadiene to cyclooctene.
- L27 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2002 ACS
- AN 1999:745198 CAPLUS
- DN 132:24789
- TI **Polymer**-protected bimetallic nanocluster **catalysts**having core/shell structure for accelerated electron transfer in
 visible-light-induced hydrogen generation
- AU Toshima, Naoki; Hirakawa, Kazutaka
- CS Department of Materials Science and Engineering, Science University of Tokyo in Yamaguchi, Onoda, 756-0884, Japan
- SO Polymer Journal (Tokyo) (1999), 31(11-2), 1127-1132 CODEN: POLJB8; ISSN: 0032-3896
- PB Society of Polymer Science, Japan
- DT Journal
- LA English
- AB A visible-light-induced electron transfer system was constructed by using tris(bipyridine)ruthenium(III), Me viologen, EDTA and colloidal metal nanocluster as photosensitizer, electron relay, sacrificial electron donor and reductive catalyst, resp. Polymer -protected Au, Pt, Pd, Rh and Ru monometallic, and Au/Pt, Au/Pd, Au/Rh and

Pt/Ru bimetallic nanocluster catalysts were prepd. as colloidal dispersions by alc.-redn., and applied to the catalysts for the above reaction. The rate of electron transfer from Me viologen cation radical to the metal nanocluster catalyst is proportional to the hydrogen generation rate at a steady state. All the electrons accepted by metal nanocluster catalysts are used for the hydrogen generation. Both electron transfer and hydrogen generation rates increase when the colloidal dispersions of bimetallic nanoclusters are used in place of the corresponding monometallic nanoclusters. An Au-core/Rh-shell structure has been suggested by UV-Vis spectrum measurement for Au/Rh bimetallic nanoclusters.

RE.CNT 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L27 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2002 ACS
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- AN 1999:529466 CAPLUS
- DN 131:162019
- TI Immobilization of palladium nanoparticles on latex supports and their potential for catalytic applications
- AU Mayer, Andrea B. R.; Mark, James E.
- CS Department Chemistry, Univ. Cincinnati, Cincinnati, OH, 45221, USA
- SO Angewandte Makromolekulare Chemie (1999), 268, 52-58 CODEN: ANMCBO; ISSN: 0003-3146
- PB Wiley-VCH Verlag GmbH
- DT Journal
- LA English
- AB Pd nanoparticles were reduced in the presence of several latex dispersions possessing different hydrophobicities. Various redn . methods were investigated, specifically the slower methods of refluxing the alc. soln. and the more rapid redn. by potassium tetrahydridoborate. In several cases the latexes showed the ability to adsorb and immobilize the Pd nanoparticles on their surface. Transmission electron microscopy was employed to show the immobilization of the metal nanoparticles on the latex surfaces, and their nanosize dimensions. The latex-metal dispersions showed catalytic activity for the hydrogenation of cyclohexene as a model reaction. A selection of water-sol. protective polymers was included to explore whether the metal nanoparticles were still immobilized. In the case of the more hydrophobic latexes, the accumulation and immobilization of the metal nanoparticles was preserved both before and after their use as hydrogenation catalysts.
- RE.CNT 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L27 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2002 ACS
- AN 1999:150983 CAPLUS
- DN 130:257756
- TI Homogeneous hydrogenation catalysis with monodisperse, dendrimer-encapsulated Pd and Pt nanoparticles
- AU Zhao, Mingqi; Crooks, Richard M.
- CS Department of Chemistry, Texas A and M University, College Station, TX, 77842-3012, USA
- No. Angewandte Chemie, International Edition (1999), 38(3), 364-366 QDI-Z5/CODEN: ACIEF5; ISSN: 1433-7851
- PB Wiley-VCH Verlag GmbH
- DT Journal
- LA English
- AB In this report we show that composite materials that consist of noble metal nanoparticles stabilized within dendrimer interiors are suitable for use as homogeneous hydrogenation catalysts. These interesting new materials are prepd. by sorbing PdII or PtII ions into hydroxyl-terminated poly(amidoamine) (PAMAM)

dendrimers (Gn-OH, were Gn represents the nth generation) where they complex with interion amine groups. Subsequent chem. redn. of the metal ions with BH4- yields dendrimer-encapsulated metal nanoparticles that contain the same no. of atoms as were preloaded into the dendrimer initially. The resulting composites are sol. in water and stable, either dry or solvated, for at least several months. Thus, the dendrimer acts as both a template for the prepn. of monodisperse nanoparticles and a porous stabilizer. Dendrimer-encapsulated Pd clusters exhibit high catalytic activity for the hydrogenation of alkenes in water. Importantly, the catalytic activity can be controlled by adjusting the size (generation) of the dendrimer; i.e., the dendrimer acts as a "nanofilter" with a synthetically controllable mesh.

RE.CNT 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 6 CAPLUS COPYRIGHT 2002 ACS

```
1997:274467 CAPLUS
AN
DN
     126:318654
TI
     Platinum nanocatalysts immobilized on latex supports
ΑU
     Mayer, Andrea B. R.; Mark, James E.
CS
     Dep. Chem. and Polymer Res. Center, Univ. Cincinnati, Cincinnati, OH,
     54221-0172, USA
     Journal of Polymer Science, Part B: Polymer Physics (1997), 35(8),
SO
     1207-1216
     CODEN: JPBPEM; ISSN: 0887-6266
PB
     Wiley
DΤ
     Journal
LA
     English
AΒ
     Several latex dispersions of different hydrophobicity were investigated
     with respect to their ability to adsorb platinum nanoparticles
     that had been reduced in their presence. Two redn. methods were
     tested, specifically the slower method of refluxing the alc.
     solns. and the more rapid method of reaction with KBH4.
     immobilization of the metal particles and their nanosize
     dimensions were demonstrated by transmission electron microscopy and their
     catalytic activity was tested by the hydrogenation of cyclohexene as a
     model reaction. Some addnl. immobilized platinum nanoparticles
     were prepd. in the presence of various protective polymers.
     This can lead to various advantages with respect to, for instance, the
     stability and the catalytic properties of these materials. Even in the
     presence of such addnl. protective polymers, the platinum nanoparticles remained immobilized for some of the hydrophobic
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=> dis hist

L27

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latexes both before and after catalytic hydrogenations.

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FILE 'CAPLUS' ENTERED AT 19:58:09 ON 09 OCT 2002
L1
        2369625 S OXID?
L2
         169395 S L1 AND AOUEOUS
L3
          24536 S L2 AND CATALYST
             67 S L3 AND NANOPARTICLE
L4
L5
              8 S L4 AND STABILIZ?
L6
              3 S L5 AND ALCOHOL
L7
              1 S L5 AND CARBOHYDRATE
L8
              2 S L5 AND PLATINUM
L9
         150451 S HYDROGENATION
L10
          14512 S L9 AND AQUEOUS
L11
           6442 S L10 AND CATALYST
L12
           1568 S L11 AND METAL
L13
              3 S L12 AND POLYMER (W) STABILIZ?
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L14	89	S	L12	AND	POLYMER
L15	5	S	L14	AND	NANOPARTICLE
L16	0	S	L15	AND	ALDEHYDE
L17	2	S	L15	AND	ALCOHOL
L18	0	S	L15	AND	CARBOHYDRATE
L19	0	S	L15	AND	FRUCTOSE
L20	891539	S	REDU	JCTI	NC
L21	68034	S	L20	AND	CATALYST
L22	15952	S	L21	AND	METAL
L23	129	S	L22	AND	NANOPARTICLE
L24	36	S	L23	AND	POLYMER
L25	0	S	L24	AND	ALDEHYDE
L26	0	S	L24	AND	CARBOHYDRATE
L27	6	S	L24	AND	ALCOHOL

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NEWS WWW

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PASSWORD:
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NEWS 2 Apr 08 "Ask CAS" for self-help around the

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Direct Dial and Telecommunication Network Access to STN

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SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST

ENTRY SESSION 0.21 0.21

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=> s amination

24906 AMINATION

249 AMINATIONS

L1 24959 AMINATION

(AMINATION OR AMINATIONS)

=> s l1 and reductive

49778 REDUCTIVE

4 REDUCTIVES

49782 REDUCTIVE

(REDUCTIVE OR REDUCTIVES)

L2 4959 L1 AND REDUCTIVE

=> s 12 and catalyst

601917 CATALYST

585163 CATALYSTS

767433 CATALYST

(CATALYST OR CATALYSTS)

L3 964 L2 AND CATALYST

=> s 13 and metal

1371087 METAL

685692 METALS

1661953 METAL

(METAL OR METALS)

L4 209 L3 AND METAL

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=> s 14 and alcohol
         165042 ALCOHOL
         124445 ALCOHOLS
         270252 ALCOHOL
                   (ALCOHOL OR ALCOHOLS)
         503294 ALC
         162712 ALCS
         587755 ALC
                   (ALC OR ALCS)
         681787 ALCOHOL
                   (ALCOHOL OR ALC)
L5
             38 L4 AND ALCOHOL
=> s 15 and aqueous
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         148294 AQUEOUS
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         933909 AQ
            119 AQS
         933982 AQ
                   (AQ OR AQS)
         968682 AQUEOUS
                   (AQUEOUS OR AQ)
              7 L5 AND AQUEOUS
L6
=> dis 16 1-7 bib abs
     ANSWER 1 OF 7 CAPLUS COPYRIGHT 2002 ACS
     2002:378142 CAPLUS
ΑN
     137:93568
DN
     Synthesis of Primary Amines: First Homogeneously Catalyzed
ΤI
     Reductive Amination with Ammonia
ΑΠ
     Gross, Thoralf; Seayad, Abdul Majeed; Ahmad, Moballigh; Beller, Matthias
CS
     Institut fuer Organische Katalyseforschung, Universitaet Rostock e.V.,
     Rostock, D-18055, Germany
     Organic Letters (2002), 4(12), 2055-2058
SO
     CODEN: ORLEF7; ISSN: 1523-7060
PB
     American Chemical Society
DТ
     Journal
T.A
     English
     The synthesis of primary amines via reductive amination
AR
     of the corresponding carbonyl compds. with aq. ammonia is achieved for the first time with sol. transition metal complexes. Up to an 86% yield and a 97% selectivity for benzylamines were
     obtained in the case of various benzaldehydes by using a Rh-
     catalyst together with water-sol. phosphine and ammonium acetate.
     In the case of aliph. aldehydes, a bimetallic catalyst based on
     Rh/Ir gave improved results.
RE.CNT 20
               THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD
               ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 2 OF 7 CAPLUS COPYRIGHT 2002 ACS
L6
     2001:122109 CAPLUS
AN
DN
     134:281120
ΤI
      .alpha.-Ketocarbonyl Peptides: A General Approach to Reactive Resin-Bound
     Intermediates in the Synthesis of Peptide Isosteres for Protease Inhibitor
     Screening on Solid Support
ΑU
     Papanikos, Alexandra; Rademann, Jorg; Meldal, Morten
     Department of Chemistry, Carlsberg Research Center, Valby, DK-2500, Den. Journal of the American Chemical Society (2001), 123(10), 2176-2181
CS
SO
     CODEN: JACSAT; ISSN: 0002-7863
```

PB

American Chemical Society

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English
LA
     CASREACT 134:281120
OS
     .alpha.-Ketocarbonyl peptides were generated from peptide precursors on
AB
     solid support via a metal-ion-catalyzed transamination. The
     reaction proceeded to completion within 2 h with glyoxylate as
     electrophile and copper(II) ions as catalyst in an aq.
     acetate buffer at pH 5.5-6.0. The variety of naturally occurring
     .alpha.-amino acid substrates gave rise to a diverse set of differentially
     functionalized ketones. The highly reactive terminal keto-carbonyls were
     prone to aldol-type dimerization and could be transferred into stable
     moieties by oxime formation, redn. to the alc., or
     reductive amination, resp. The .alpha.-keto-carbonyl
     peptides were efficient in nucleophilic addn. of C-nucleophiles such as
     phosphono-ylides and allylsilanes.
              THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 20
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 3 OF 7 CAPLUS COPYRIGHT 2002 ACS
L6
     2000:666737 CAPLUS
AN
DN
     133:254142
ΤI
     Catalytic method for modifying carbohydrates, alcohols,
     aldehydes or polyhydroxy compounds
     Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter;
IN
     Haji Begli, Alireza
     Sudzucker Aktiengesellschaft, Germany
PA
SO
     PCT Int. Appl., 45 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     German
FAN.CNT 1
     PATENT NO.
                     KIND DATE
                                          APPLICATION NO.
                                                           DATE
     ______
                     ____
                           _____
                                          _____
     WO 2000055165 A1
                                          WO 2000-EP2351
PΤ
                           20000921
                                                           20000316
        W: AU, CA, IL, US
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
                                          DE 1999-19911504 19990316
                           20001019
     DE 19911504
                      A1
                     A1
                                          EP 2000-925117
     EP 1165580
                           20020102
                                                           20000316
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             IE, FI
     AU 747812
                                          AU 2000-43953
                      B2
                           20020523
                                                           20000316
PRAI DE 1999-19911504 A
                           19990316
     WO 2000-EP2351
                      W
                           20000316
AB
     Industrial conversion of the title compds. in aq. phase is
     carried out in the presence of metal catalysts
     consisting of polymer-stabilized nanoparticles. A catalyst of
     this type is not deactivated by the conversion reaction as long as the
     stabilizing interaction between the polymer and the nanoparticles is
     maintained. For example, activity of an Al2O3-supported,
     poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn.
     given) in oxidn. of sorbose with O remained unchanged after 10 repeated
     expts. whereas the activity of a customary Al2O3-supported Pt
     catalyst decreased to .apprx.35% after 10 runs.
             THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 6
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 4 OF 7 CAPLUS COPYRIGHT 2002 ACS
L6
    1998:693414 CAPLUS
ΑN
DN
    129:275633
TΤ
    Metal catalysts and preparation of the
    catalysts and (N-substituted) amines
    Muraishi, Teruo; Kato, Kozo
ΙN
PΑ
    Mitsui Chemicals Inc., Japan
```

DT

Journal

SO Jpn. Kokai Tokkyo Koho, 7 pp. CODEN: JKXXAF DTPatent Japanese LAFAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE JP 10287628 A2 19981027 JP 1997-89526 19970408 PΙ (N-substituted) amines are prepd. by treatment of NH3, primary amines, or AB secondary amines with (i) alcs. or (ii) aldehydes and H in the presence of catalysts prepd. by redn. of malachite- and Al(OH)3-free precursors contg. Cu, Al, and Cr, Mn, Fe, Co, Ni, and/or Zn. Aq. soln. contq. Cu(NO3)2, Ni(NO3)2, and Al(NO3)3 was treated with aq. Na2CO3 to give hydrotalcite-like substance, which was reduced in lauryl alc. NHMe2 and H were passed through the catalyst-contg. reactor at 200.degree. over 10 h to give colorless products contg. 91.8% lauryldimethylamine. ANSWER 5 OF 7 CAPLUS COPYRIGHT 2002 ACS L6 1995:401307 CAPLUS AΝ DN 122:160152 TΙ Process for producing amines by reductive amination in the presence of a cobalt catalyst. Furutani, Atsushi; Hibi, Takuo; Yamamoto, Michio; Tanaka, Kazuyuki; Tada, IN Kazuhiro; Fukao, Masami; Suzukamo, Gohfu PA Sumitomo Chemical Co., Ltd., Japan Eur. Pat. Appl., 13 pp. SO CODEN: EPXXDW DTPatent English LAFAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE EP 623585 A1 19941109 EP 1994-300903 19940208 EP 623585 B1 19980422 PΤ R: BE, DE, FR, GB JP 07101910 A2 19950418 JP 1994-6466 19940125
CA 2115024 AA 19941028 CA 1994-2115024 19940204
US 5589596 A 19961231 US 1994-194328 19940208
PRAI JP 1993-101074 19930427
JP 1993-180248 19930721
JP 1993-180249 19930721 JP 1993-196041 19930806 JP 1993-197339 19930809 OS CASREACT 122:160152 AB A process is disclosed for producing amines by reductive amination of cyclic ketones or their imino derivs., characterized by use of a cobalt catalyst contg. an alk. earth metal carbonate and/or lanthanum oxide. The new catalysts give high yields, are highly active, and are usable on a com. scale. For example, an aq. soln. of Co and Cu nitrates was treated with Ca carbonate, heated to 80.degree., and treated with aq. Na carbonate to give a ppt., which was retreated with aq. Na carbonate, dried, heated in N at 320.degree., cooled, granulated, and hydrogenated at 280.degree. to give a catalyst. 3-Cyano-3,5,5-trimethylcyclohexanone was then passed with MeOH and liq. NH3 through a first reactor contg. active C at 24.degree. and 150 kg/cm2G to give the imine deriv. in 97.7% yield. This was passed through the above catalyst in a second reactor at 121.degree. and the same pressure to give 3-aminomethyl-3,5,5-trimethylcyclohexylamine (I) in 99.4% yield, plus minor amts. of 2 byproducts. A comparison catalyst without the Cu nitrate or the Ca carbonate gave only 90.7% yield of I in the second step, with 5.8% 3-aminomethyl-3,5,5-trimethylcyclohexyl

alc. and 3.2% 1,3,3-trimethyl-6-azabicyclo[3.2.1]octane as

byproducts.

```
ANSWER 6 OF 7 CAPLUS COPYRIGHT 2002 ACS
L6
AN
     1993:603755 CAPLUS
     119:203755
DN
     Process for the production of aminopolyols
ΤI
     Beck, Roland Herwig; Kalff, Norbert Johannes; Roeper, Harald W. W.
IN
     Cerestar Holding BV, Neth.
PΑ
     Eur. Pat. Appl., 7 pp.
SO
     CODEN: EPXXDW
DT
     Patent
     English
LA
FAN.CNT 1
                                           APPLICATION NO. DATE
     PATENT NO. KIND DATE
                      ____
                                             -----
     EP 536939 A1 19930414
                                            EP 1992-308919 19920930
PΙ
                       B1 19960320
     EP 536939
     EP 536939 B2 19990310
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, NL, PT, SE
     AT 135680 E 19960415
ES 2084947 T3 19960516
                                           AT 1992-308919 19920930
                                             ES 1992-308919
                                                                19920930
PRAI GB 1991-21279
                             19911008
     CASREACT 119:203755
OS
     The title process comprises reacting a monosaccharide or a reducing
ΑB
     oligosaccharide with NH3 or an aliph. amine having a replaceable amino
     hydrogen atom in the presence of H2 and a base metal
     catalyst and is characterized by the addn. to the reaction medium
     of a compd. which, under the reaction conditions is capable of providing
     NH4+ in the reaction medium and is not, or does not give rise to, a poison
     for the base metal catalyst. Thus, a mixt. of
     aq. glucose and NH4OAc was hydrogenated over Raney Co with the
     addn. of liquefied NH3 at 5 to 100 bar and -40 to 95.degree. to give 83.4
     wt.% 1-aminosorbitol, 3.1 wt.% 2-aminosorbitol, 3.5 wt.% 2-aminomannitol,
     2.2 wt.% sorbitol, and 2.6 wt.% disorbitylamine.
     ANSWER 7 OF 7 CAPLUS COPYRIGHT 2002 ACS
L6
AN
     1987:5981 CAPLUS
DN
     106:5981
ΤI
     2,2,6,6-Tetraalkyl-4-piperidylamines
     Malz, Russell E., Jr.; Greenfield, Harold Uniroyal Chemical Co., Inc., USA
ΙN
PΑ
SO
     U.S., 5 pp.
     CODEN: USXXAM
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
                      KIND DATE
                                    APPLICATION NO.
                                                               DATE
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                                             -----
                                                               -----
                A 19860819
A 19921128
A 19870225
A 19870317
A1 19870114
B1 19910130
PΙ
     US 4607104
                                             US 1985-754378
                                                                19850711
     IN 171623
                                             IN 1986-DE494
                                                                19860604
     ZA 8604298
                                             ZA 1986-4298
                                                                19860609
     BR 8602855
                                             BR 1986-2855
                                                                19860619
     EP 208455
                                             EP 1986-304837
                                                                19860624
     EP 208455
         R: AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE
     AT 60583
                 E 19910215
                                             AT 1986-304837
                                                                19860624
    CA 1283108 A1 19910416

CN 86104693 A 19870225

CN 1005976 B 19891206

AU 8660042 A1 19870115

AU 589196 B2 19891005

JP 62030757 A2 19870209

JP 05002669 B4 19930113

ES 2000661 A6 19880316
                                             CA 1986-513288
                                                                19860708
                                             CN 1986-104693
                                                                19860709
                                             AU 1986-60042
                                                                19860710
                                             JP 1986-162917
                                                                19860710
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ES 1986-246

19860711

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PRAI US 1985-754378 19850711
EP 1986-304837 19860624
OS CASREACT 106:5981
GI
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R^2 \\
R^3 \\
N-R^6 \\
R^5 \\
R^4
\end{array}$

The title compds. I (R1 = C1-18 alkyl, C5-6 cycloalkyl, C7-9 aralkyl; R2-5 = C1-8 alkyl; R6 = H, OH, C1-8 alkyl, alkoxy, C2-10 alkylcarbonyl, arylcarbonyl; n = 1-4), useful as UV stabilizers, are prepd. by reductive amination over pyrophoric catalysts (Pt, Ni or Co) with reduced fire hazard by using H2O or aq. alcs. as solvents. Thus, 34.8 g H2N(CH2)6NH2, 97.65 g 2,2,6,6-tetramethyl-4-piperidone, 120 mL water, and 1.5 g 5% Pt/C were heated at 80.degree./600-800 psig with H to give 89% N,N'-bis(2,2,6,6-tetramethyl-4-piperidyl)hexamethylenediamine, vs. 93% with MeOH instead of H2O.

=> dis hist

(FILE 'HOME' ENTERED AT 20:13:40 ON 09 OCT 2002)

FILE 'CAPLUS' ENTERED AT 20:13:50 ON 09 OCT 2002 L124959 S AMINATION 4959 S L1 AND REDUCTIVE L2 L3 964 S L2 AND CATALYST 209 S L3 AND METAL L4L5 38 S L4 AND ALCOHOL L6 7 S L5 AND AQUEOUS L7 0 S L6 AND PROMOTER => s 14 and aldehyde 89500 ALDEHYDE

139257 ALDEHYDE
(ALDEHYDE OR ALDEHYDES)
L8 35 L4 AND ALDEHYDE

84224 ALDEHYDES

=> s 18 and aqueous
148293 AQUEOUS
1 AQUEOUSES
148294 AQUEOUS
(AQUEOUS OR AQUEOUSES)
933909 AQ
119 AQS

933982 AQ

L9 6 L8 AND AQUEOUS

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=> dis 19 1-9 bib abs
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ANSWER 1 OF 6 CAPLUS COPYRIGHT 2002 ACS 1.9

2002:378142 CAPLUS ΑN

137:93568 DN

Synthesis of Primary Amines: First Homogeneously Catalyzed TΙ Reductive Amination with Ammonia

Gross, Thoralf; Seayad, Abdul Majeed; Ahmad, Moballigh; Beller, Matthias ΑU

Institut fuer Organische Katalyseforschung, Universitaet Rostock e.V., CS

Rostock, D-18055, Germany Organic Letters (2002), 4(12), 2055-2058 Acade - not crowd CODEN: ORLEF7; ISSN: 1523-7060 SO

PΒ American Chemical Society

DTJournal

LA English

AR The synthesis of primary amines via reductive amination of the corresponding carbonyl compds. with aq. ammonia is achieved for the first time with sol. transition metal complexes. Up to an 86% yield and a 97% selectivity for benzylamines were obtained in the case of various benzaldehydes by using a Rhcatalyst together with water-sol. phosphine and ammonium acetate. In the case of aliph. aldehydes, a bimetallic catalyst based on Rh/Ir gave improved results.

THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 20 ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 6 CAPLUS COPYRIGHT 2002 ACS L9

2000:666737 CAPLUS ΑN

133:254142 DN

TΙ Catalytic method for modifying carbohydrates, alcohols, aldehydes or polyhydroxy compounds

Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter; IN Haji Begli, Alireza

immertant Sudzucker Aktiengesellschaft, Germany PΑ

PCT Int. Appl., 45 pp. SO CODEN: PIXXD2

DTPatent

LA German

FAN.CNT 1

PATENT NO. KIND DATE APPLICATION NO. DATE WO 2000055165 PΤ A1 20000921 WO 2000-EP2351 20000316 W: AU, CA, IL, US RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE DE 19911504 20001019 DE 1999-19911504 19990316 A1 EP 2000-925117 EP 1165580 A1 20020102 20000316 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI AU 747812 B2 20020523 AU 2000-43953 20000316 PRAI DE 1999-19911504 A 19990316

WO 2000-EP2351 W 20000316 Industrial conversion of the title compds. in aq. phase is AB

carried out in the presence of metal catalysts consisting of polymer-stabilized nanoparticles. A catalyst of this type is not deactivated by the conversion reaction as long as the stabilizing interaction between the polymer and the nanoparticles is maintained. For example, activity of an Al203-supported, poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn.

given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs.

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
L9 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2002 ACS
```

AN 1998:693414 CAPLUS

DN 129:275633

TI Metal catalysts and preparation of the catalysts and (N-substituted) amines

IN Muraishi, Teruo; Kato, Kozo

PA Mitsui Chemicals Inc., Japan

SO Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1
PATENT NO. KIND DATE APPLICATION NO. DATE

| ΡI | JP 10287628 | A2 | 19981027 | JP 1997- | 89526 | 19970408 | |
|----|------------------|----------|-------------|---------------|----------|------------|-------------|
| AB | (N-substituted) | | | | | | amines, or |
| | secondary amines | | | | | | |
| | presence of cata | alysts p | prepd. by r | edn. of malac | hite- an | d | |
| | Al(OH)3-free pre | ecursors | s contg. Cu | , Al, and Cr, | Mn, Fe, | Co, Ni, a | and/or Zn. |
| | Aq. soln. contg. | . Cu (NO | 3)2, Ni(NO3 |)2, and Al(NO | 3)3 was | treated wi | th |
| | aq. Na2CO3 to gi | ive hydi | rotalcite-l | ike substance | , which | was reduce | ed |
| | in lauryl alc. | NHMe2 a | and H were | passed throug | h the ca | talyst | |
| | -contg. reactor | at 200. | degree. ov | er 10 h to gi | ve color | less produ | icts contg. |
| | 91.8% lauryldime | ethylami | ine. | _ | | _ | _ |

- L9 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2002 ACS
- AN 1991:535511 CAPLUS
- DN 115:135511
- TI Process for the preparation of N, N-dimethylamines
- IN Kampmann, Detlef; Kniep, Claus; Lukas, Rainer
- PA Hoechst A.-G., Germany
- SO Ger. Offen., 6 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

| | PATENT NO. | | KIND | DATE | | APPLICATION NO. | DATE |
|------|-------------------------|-------|--------------|-----------------------|-----|-----------------------------------|----------|
| PI | DE 3942793
EP 435072 | | A1 | 19910627
19910703 | | DE 1989-3942793
EP 1990-123912 | 19891223 |
| | EP 435072 | | A2
A3 | 19920304 | | EP 1990-123912 | 19901212 |
| | EP 435072
R: AT | , BE, | B1
CH, DE | 19940427
, ES, FR, | GB, | IT, LI, NL, SE | |
| | AT 104950 | | E | 19940515 | | AT 1990-123912 | 19901212 |
| | ES 2055855 | | Т3 | 19940901 | | ES 1990-123912 | 19901212 |
| | CA 2032362 | | AA | 19910624 | | CA 1990-2032362 | 19901214 |
| | JP 0621999 | - | A2 | 19940809 | | JP 1990-402867 | 19901217 |
| | JP 0707215 | - | B4 | 19950802 | | | |
| | AU 9068373 | | A1 | 19910627 | | AU 1990-68373 | 19901221 |
| | AU 634007 | | B2 | 19930211 | | | |
| PRAI | DE 1989-39 | | | 19891223 | | | |
| | EP 1990-12 | 3912 | | 19901212 | | | |

AB The continuous prepn. of N,N-dimethylamines by the reaction of aldehydes with Me2NH and hydrogen under pressure at high temp. in the presence of Ni, Co, Cu, Mn, Fe, Rh, Pd and/or Pt-contg. hydrogenation catalysts is claimed. After remaining starting material (Me2NH) and hydrogen are removed, 0.1-25% by wt. HCHO or HCHO-forming substance are added and the mixt. is distd. This process permits nearly complete

removal of secondary N-methylamines which are formed as by products. A reactor contg. 300 mL catalyst RCH Ni52/35 (tablets; Ni catalyst on kieselguhr) was filled with Me2NBu and then charged with PrCHO (65 mL) and Me2NH (200 mL) at 105-110.degree. and 8 MPa and hydrogen was charged at 34 L/h; remaining hydrogen and Me2NH were removed and during the subsequent distn. 37% aq. HCHO (.apprx.3% with respect to Me2NBu) was fed into the crude product mixt. at the bottom of the column. The distillate contained 99.65% by wt. Me2NBu and 0.02% by wt. MeNHBu. Omission of feed of aq. HCHO gave a distillate contg. 98.09% by wt. Me2NBu and 1.22% by wt. MeNHBu.

- L9 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2002 ACS
- AN 1982:616038 CAPLUS
- DN 97:216038
- TI Araliphatic aldehydes and/or amines
- IN Gramlich, Walter; Heilen, Gerd; Mercker, Hans Jochen; Siegel, Hardo
- PA BASF A.-G., Fed. Rep. Ger.
- SO Ger. Offen., 13 pp.

CODEN: GWXXBX

DT Patent

LA German

FAN.CNT 1

| | PA' | FENT NO. | KIND | DATE | APPLICATION N | O. DATE | | | |
|----|-------|--------------|--------|----------|---------------|-------------|--|--|--|
| | | | | | | | | | |
| ΡI | DE | 3105446 | A1 | 19820902 | DE 1981-31054 | 46 19810214 | | | |
| | EP | 58326 | A1 | 19820825 | EP 1982-10067 | 0 19820130 | | | |
| | EP | 58326 | B1 | 19840606 | | | | | |
| | | R: CH, DE, | FR, GB | , NL | | | | | |
| | JP | 57163334 | A2 | 19821007 | JP 1982-19006 | 19820210 | | | |
| PR | AI DE | 1981-3105446 | | 19810214 | | | | | |
| GΙ | | | | | | | | | |

- AB I (R, R1, R2 = H, C1-8 alkyl, C4-7 cycloalkyl, C1-5 alkoxy, OH, tertiary amino, etc.; R3 = H, C1-10 alkyl, etc.; X = CHO or NR4R5; R4, R5 = C1-10 alkyl or R4R5N = 5-7-membered heterocycle) were prepd. by aldol condensation and hydrogenation or reductive amination.

 Thus, 4-Me3CC6H4CHO, EtCHO, MeOH, and aq. NaOH first at 30-40.degree. under N, then at 110.degree. under 2 bar H (metal oxide catalyst) gave 4-Me3CC6H4CH2CHMeCHO. Similarly, but with morpholine or hexamethylenimine present in the 2nd step, the reaction gave II or III.
- L9 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2002 ACS
- AN 1982:199072 CAPLUS
- DN 96:199072
- TI N-Alkylalkylenediamines
- IN Tahara, Susumu; Nishihira, Keigo; Miyatake, Takashi; Sawada, Hiroyuki; Kita, Junichiro

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PA
     Ube Industries, Ltd., Japan
SO
     Ger. Offen., 13 pp.
     CODEN: GWXXBX
DT
     Patent
     German
T.A
FAN.CNT 1
                                        APPLICATION NO. DATE
     PATENT NO.
                    KIND DATE
     DE 3128810
                     A1 19820318
                                          DE 1981-3128810 19810721
PΤ
     JP 57035545
                     A2 19820226
                                          JP 1980-111590
                                                          19800813
     JP 60033423
                     B4
                          19850802
     US 4373107
                     Α
                          19830208
                                          US 1981-286030
                                                           19810722
                           19820303
     GB 2082172
                                          GB 1981-22958
                                                          19810724
                      А
                          19840510
     GB 2082172
                      B2
PRAI JP 1980-111590
                           19800813
                                                                         not aqueous
     N-Alkylalkylenediamines were prepd. by gradual addn. of an aliph.
     aldehyde to an alkylenediamine in an org. solvent over a Pt group
     metal in an atm. of H under pressure. Thus, a mixt. of 37 g 90%
     aq. AcH and 100 mL MeOH was added over 8 h to 45 g H2NCH2CH2NH2
     (I), 2 g 5% Pd-C, and 100 mL MeOH in a 0.5 L autoclave at 100.degree./7
     bar H overpressure, and the mixt. heated as above 4 h to give 40.9%
     conversion of I with 78.9% selectivity to EtNHCH2CH2NH2.
=> s 14 and carbohydrate
        108745 CARBOHYDRATE
        116172 CARBOHYDRATES
        176447 CARBOHYDRATE
                 (CARBOHYDRATE OR CARBOHYDRATES)
            2 L4 AND CARBOHYDRATE
L10
=> dis 110 1-2 bib abs
L10 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2002 ACS
     2000:666737 CAPLUS
AN
DN
     133:254142
ΤI
    Catalytic method for modifying carbohydrates, alcohols,
     aldehydes or polyhydroxy compounds
    Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter;
ΙN
    Haji Begli, Alireza
PA
    Sudzucker Aktiengesellschaft, Germany
    PCT Int. Appl., 45 pp.
SO
    CODEN: PIXXD2
DΤ
    Patent
LA
    German
FAN.CNT 1
    PATENT NO.
                     KIND DATE
                                        APPLICATION NO.
                                                           DATE
                     ____
                           _____
                                          -----
                                                           -----
    WO 2000055165
                    A1
                           20000921
                                          WO 2000-EP2351
PΤ
                                                           20000316
        W: AU, CA, IL, US
        RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, SE
    DE 19911504
                           20001019
                                          DE 1999-19911504 19990316
                      Α1
    EP 1165580
                                          EP 2000-925117
                           20020102
                      A1
                                                           20000316
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, FI
    AU 747812
                      B2
                           20020523
                                          AU 2000-43953
                                                           20000316
PRAI DE 1999-19911504 A
                           19990316
    WO 2000-EP2351 W
                           20000316
AB
    Industrial conversion of the title compds. in ag. phase is carried out in
    the presence of metal catalysts consisting of
    polymer-stabilized nanoparticles. A catalyst of this type is
    not deactivated by the conversion reaction as long as the stabilizing
```

interaction between the polymer and the nanoparticles is maintained. For

example, activity of an Al2O3-supported, poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn. given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs.

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
L10 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2002 ACS
     1997:530000 CAPLUS
AN
     127:150351
DN
ΤI
    Metal-catalyzed oxidation and reduction of carbohydrates
ΑIJ
     Abbadi, A.; Van Bekkum, H.
     Laboratory of Organic Chemistry and Catalysis, Delft University of
CS
     Technology, Delft, 2628 BL, Neth.
     Carbohydrates as Organic Raw Materials III, developed from a Workshop,
SO
     Wageningen, Nov. 28-29, 1994 (1996), Meeting Date 1994, 37-65. Editor(s):
     Van Bekkum, Herman; Roeper, Harald; Voragen, Fons. Publisher: VCH,
     Weinheim, Germany.
     CODEN: 64UXAZ
DT
```

Conference; General Review

LA English

AB A review with 210 refs. on metal-catalyzed oxidn. and hydrogenation of carbohydrates is presented. Bi-promoted Pd is presently the catalyst of choice for the oxidn. of aldoses to aldonic acids. Selective oxidn. at C-2 of aldonic acids to 2-keto acids can be achieved by applying similarly promoted Pt in weakly acidic medium. The noble metal-catalyzed oxidn. of primary alc. groups still needs improvement; best results are obtained in a continuous process using supported Pt and operating at low O concn. Carbonyl group hydrogenation is generally performed over Ni catalysts, but Ru catalysts are also used. Mainly Ni and Pt catalysts are applied for reductive amination of carbohydrates. Finally, carbohydrate hydrogenolysis under severe conditions to give glycerol and 1,2-propanediol is discussed.

=> dis hist

(FILE 'HOME' ENTERED AT 20:13:40 ON 09 OCT 2002)

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FILE 'CAPLUS' ENTERED AT 20:13:50 ON 09 OCT 2002
L1
          24959 S AMINATION
L2
           4959 S L1 AND REDUCTIVE
L3
            964 S L2 AND CATALYST
L4
            209 S L3 AND METAL
L5
             38 S L4 AND ALCOHOL
              7 S L5 AND AQUEOUS
L6
L7
              0 S L6 AND PROMOTER
\Gamma8
             35 S L4 AND ALDEHYDE
L9
              6 S L8 AND AQUEOUS
              2 S L4 AND CARBOHYDRATE
L10
=> s 16 and electrodialysis
          9460 ELECTRODIALYSIS
            17 ELECTRODIALYSES
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                  (ELECTRODIALYSIS OR ELECTRODIALYSES)
L11
             O L6 AND ELECTRODIALYSIS
=> s 19 and electrodialysis
          9460 ELECTRODIALYSIS
            17 ELECTRODIALYSES
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9465 ELECTRODIALYSIS

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(ELECTRODIALYSIS OR ELECTRODIALYSES)
0 L9 AND ELECTRODIALYSIS
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=> s 14 and nanoparticle 11088 NANOPARTICLE 17675 NANOPARTICLES 18714 NANOPARTICLE (NANOPARTICLE OR NANOPARTICLES) L13 1 L4 AND NANOPARTICLE => dis 113 bib abs

L13 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2002 ACS

2000:666737 CAPLUS AN

133:254142 DN

L12

Catalytic method for modifying carbohydrates, alcohols, aldehydes or TΙ polyhydroxy compounds

Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter; TN Haji Begli, Alireza

PA Sudzucker Aktiengesellschaft, Germany

PCT Int. Appl., 45 pp. SO

CODEN: PIXXD2

DΤ Patent

LA German

FAN.CNT 1

APPLICATION NO. DATE PATENT NO. KIND DATE _____ ------WO 2000055165 A1 20000921 WO 2000-EP2351 20000316 PT W: AU, CA, IL, US RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE 20001019 DE 1999-19911504 19990316 DE 19911504 A1 EP 1165580 20020102 EP 2000-925117 20000316 A1 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI AU 747812 20020523 AU 2000-43953 20000316 В2 PRAI DE 1999-19911504 A 19990316

WO 2000-EP2351 W 20000316

Industrial conversion of the title compds. in aq. phase is carried out in AB the presence of metal catalysts consisting of polymer-stabilized nanoparticles. A catalyst of this type is not deactivated by the conversion reaction as long as the stabilizing interaction between the polymer and the nanoparticles is maintained. For example, activity of an Al2O3-supported, poly(vinylpyrrolidone)-stabilized Pt colloid catalyst (prepn. given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs.

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 6 ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 14 and alloy 564477 ALLOY 443476 ALLOYS 720959 ALLOY (ALLOY OR ALLOYS) L14 5 L4 AND ALLOY

=> s 114 and alcohol 165042 ALCOHOL 124445 ALCOHOLS 270252 ALCOHOL (ALCOHOL OR ALCOHOLS)

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503294 ALC
        162712 ALCS
        587755 ALC
                  (ALC OR ALCS)
        681787 ALCOHOL
                  (ALCOHOL OR ALC)
L15
             0 L14 AND ALCOHOL
=> s metal
       1371087 METAL
        685692 METALS
       1661953 METAL
L16
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=> s 116 and catalyst
        601917 CATALYST
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L17
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        708926 POLYMERS
       1158743 POLYMER
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L19
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        626498 OXIDN
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L22
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=> dis 121 bib abs
L21 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2002 ACS
AN
     2002:720926 CAPLUS
     137:222515
DN
ΤI
     Catalysis by gold nanoparticles
     Grisel, Ruud; Weststrate, Kees-Jan; Gluhoi, Andrea; Nieuwenhuys, Bernard
ΑU
     Ε.
```

```
Leiden Institute of Chemistry, Leiden University, Leiden, 2300, Neth.
CS
     Gold Bulletin (London, United Kingdom) (2002), 35(2), 39-45
SO
     CODEN: GOBUFW; ISSN: 1027-8591
     World Gold Council
PR
     Journal; General Review
DT
LA
     English
     A review; gold catalysts have superior activity in CO and other
AB
     oxidns. at low temps. Both a small (.apprx. 5nm) particle size
     and the presence of a partly reducible oxide (ceria or a transition
     metal oxide) have a beneficial effect on the catalyst
     performance. The present paper reviews our recent studies focused on
     understanding the specific role of the Au particle size and that of the
     oxide (MO). Our personal viewpoint on gold catalysis is outlined. The
     effects of Au particle size and of the oxidic additive are distinguished
     by using several alumina-supported gold catalysts having
     different gold particle sizes and various oxidic additives. The most
     active catalyst in CO oxidn. is the multicomponent
     catalyst Au/MgO/MnOx/Al2O3 with MgO being a stabilizer
     for the Au particle size and MnOx being the cocatalyst.
     catalyst also exhibits good performance in selective oxidn
     . of CO in a hydrogen atm., a reaction relevant for the development of
     polymer electrolyte fuel cell technol.
=> s 121 and alcohol
        165042 ALCOHOL
        124445 ALCOHOLS
        270252 ALCOHOL
                 (ALCOHOL OR ALCOHOLS)
        503294 ALC
        162712 ALCS
        587755 ALC
                 (ALC OR ALCS)
        681787 ALCOHOL
                 (ALCOHOL OR ALC)
L23
             1 L21 AND ALCOHOL
=> dis 123 bib abs
     ANSWER 1 OF 1 CAPLUS COPYRIGHT 2002 ACS
ΑN
     2000:666737 CAPLUS
DN
     133:254142
ΤI
     Catalytic method for modifying carbohydrates, alcohols,
     aldehydes or polyhydroxy compounds
ΙN
     Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter;
     Haji Begli, Alireza
PΑ
     Sudzucker Aktiengesellschaft, Germany
SO
     PCT Int. Appl., 45 pp.
     CODEN: PIXXD2
DТ
     Patent
LA
     German
FAN.CNT 1
     PATENT NO.
                      KIND
                            DATE
                                           APPLICATION NO.
                                                            DATE
PΙ
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                                           WO 2000-EP2351
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             PT, SE
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                                           DE 1999-19911504 19990316
                       A1
                            20020102
                                           EP 2000-925117
     EP 1165580
                       Α1
                                                            20000316
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
     AU 747812
                       B2
                            20020523
                                           AU 2000-43953
                                                             20000316
PRAI DE 1999-19911504 A
                            19990316
```

```
WO 2000-EP2351
                      W
                            20000316
     Industrial conversion of the title compds. in aq. phase is carried out in
AB
     the presence of metal catalysts consisting of
     polymer-stabilized nanoparticles. A
     catalyst of this type is not deactivated by the conversion
     reaction as long as the stabilizing interaction between the
     polymer and the nanoparticles is maintained. For
     example, activity of an Al203-supported, poly(vinylpyrrolidone)-
     stabilized Pt colloid catalyst (prepn. given) in
     oxidn. of sorbose with O remained unchanged after 10 repeated
     expts. whereas the activity of a customary Al2O3-supported Pt
     catalyst decreased to .apprx.35% after 10 runs.
              THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 6
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> s 121 and carbohydrate
        108745 CARBOHYDRATE
        116172 CARBOHYDRATES
        176447 CARBOHYDRATE
                 (CARBOHYDRATE OR CARBOHYDRATES)
             1 L21 AND CARBOHYDRATE
L24
=> s 121 and fructose
         53219 FRUCTOSE
            82 FRUCTOSES
         53232 FRUCTOSE
                 (FRUCTOSE OR FRUCTOSES)
L25
             1 L21 AND FRUCTOSE
=> dis 125 bib abs
    ANSWER 1 OF 1 CAPLUS COPYRIGHT 2002 ACS
     2000:666737 CAPLUS
ΑN
DN
     133:254142
TΤ
     Catalytic method for modifying carbohydrates, alcohols, aldehydes or
     polyhydroxy compounds
     Capan, Emine; Hahnlein, Marc Sascha; Prusse, Ulf; Vorlop, Klaus-Dieter;
ΙN
     Haji Begli, Alireza
PΑ
     Sudzucker Aktiengesellschaft, Germany
SO
     PCT Int. Appl., 45 pp.
     CODEN: PIXXD2
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     Patent
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FAN.CNT 1
     PATENT NO.
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                                          WO 2000-EP2351
     WO 2000055165
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        W: AU, CA, IL, US
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
             PT, SE
     DE 19911504
                      Α1
                            20001019
                                           DE 1999-19911504 19990316
                     A1
     EP 1165580
                                          EP 2000-925117 20000316
                            20020102
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
     AU 747812
                           20020523
                                          AU 2000-43953
                                                           20000316
                      B2
PRAI DE 1999-19911504 A
                           19990316
     WO 2000-EP2351
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                           20000316
AB
     Industrial conversion of the title compds. in aq. phase is carried out in
     the presence of metal catalysts consisting of
     polymer-stabilized nanoparticles. A
     catalyst of this type is not deactivated by the conversion
     reaction as long as the stabilizing interaction between the
    polymer and the nanoparticles is maintained. For
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stabilized Pt colloid catalyst (prepn. given) in oxidn. of sorbose with O remained unchanged after 10 repeated expts. whereas the activity of a customary Al2O3-supported Pt catalyst decreased to .apprx.35% after 10 runs. RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT => dis hist (FILE 'HOME' ENTERED AT 20:13:40 ON 09 OCT 2002) FILE 'CAPLUS' ENTERED AT 20:13:50 ON 09 OCT 2002 24959 S AMINATION L1L2 4959 S L1 AND REDUCTIVE L3 964 S L2 AND CATALYST 209 S L3 AND METAL L4L5 38 S L4 AND ALCOHOL 7 S L5 AND AQUEOUS L6 0 S L6 AND PROMOTER L7 35 S L4 AND ALDEHYDE L8 L9 6 S L8 AND AQUEOUS 2 S L4 AND CARBOHYDRATE L10 0 S L6 AND ELECTRODIALYSIS L11 L12 0 S L9 AND ELECTRODIALYSIS L13 1 S L4 AND NANOPARTICLE L14 5 S L4 AND ALLOY L15 0 S L14 AND ALCOHOL 1661953 S METAL L16 L17 140657 S L16 AND CATALYST L18 593 S L17 AND NANOPARTICLE L19 108 S L18 AND POLYMER T₁2.0 30 S L19 AND STABILIZ? L21 7 S L20 AND OXIDAT? L22 1 S L21 AND ALDEHYDE L23 1 S L21 AND ALCOHOL L24 1 S L21 AND CARBOHYDRATE L25 1 S L21 AND FRUCTOSE => s 120 and hydrogenation or reduction 150203 HYDROGENATION 1908 HYDROGENATIONS 150451 HYDROGENATION (HYDROGENATION OR HYDROGENATIONS) 262772 REDUCTION 5081 REDUCTIONS 265419 REDUCTION (REDUCTION OR REDUCTIONS) 742417 REDN 39209 REDNS 766916 REDN (REDN OR REDNS) 891539 REDUCTION (REDUCTION OR REDN) L26 891546 L20 AND HYDROGENATION OR REDUCTION => s 120 and hydrogenation 150203 HYDROGENATION 1908 HYDROGENATIONS 150451 HYDROGENATION (HYDROGENATION OR HYDROGENATIONS)

19 L20 AND HYDROGENATION

L27

example, activity of an Al2O3-supported, poly(vinylpyrrolidone)-

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=> s 127 and aldehyde
         89500 ALDEHYDE
         84224 ALDEHYDES
        139257 ALDEHYDE
                  (ALDEHYDE OR ALDEHYDES)
             O L27 AND ALDEHYDE
L28
=> s 127 and alcohol
        165042 ALCOHOL
        124445 ALCOHOLS
        270252 ALCOHOL
                  (ALCOHOL OR ALCOHOLS)
        503294 ALC
        162712 ALCS
        587755 ALC
                  (ALC OR ALCS)
        681787 ALCOHOL
                  (ALCOHOL OR ALC)
L29
             3 L27 AND ALCOHOL
=> dis 129 1-3 bib abs
     ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS
ΑN
     2001:65241 CAPLUS
     134:268058
DN
TΤ
     Synthesis and functionalities of poly(N-vinylalkyloamide). XIII.
     Temperature and pH dependence of the catalytic activity of colloidal
     platinum nanoparticles stabilized by
     poly[(vinylamine)-co-(N-vinylisobutyramide)]
     Chen, Chun-Wei; Arai, Kumiko; Yamamoto, Kazuya; Serizawa, Takeshi; Akashi,
ΑU
     Mitsuru
     Dep. Applied Chem. Chemical Eng., Kagoshima Univ., Kagoshima, 890-0065,
CS
     Japan
     Macromolecular Chemistry and Physics (2000), 201(18), 2811-2819
SO
     CODEN: MCHPES; ISSN: 1022-1352
PB
     Wiley-VCH Verlag GmbH
DT
     Journal
     Enalish
LA
     Colloidal platinum nanoparticles in the size range of 5-35 .ANG.
AB
     have been successfully prepd. in water at room temp. by NaBH4 redn. of
     ionic platinum in the presence of poly[(vinylamine)-co-(N-
     vinylisobutyramide)] (PVAm-co-PNVIBA). The temp.- and pH-responsive
     copolymer was used for the first time as the stabilizer of
     colloidal metal particles. Three PVAm-co-PNVIBA copolymers with PVAm contents of 4.1, 8.3, and 19.8 mol-% were examd. The particle size
     and morphol. of the platinum colloids varied with the copolymer compn., as
     confirmed by TEM measurements. The polymer-stabilized
     Pt nanoparticles pptd. on heating above their crit. flocculation
     temps. (CFTs), which were strongly dependent on the soln. pH and the
     copolymer compn. The CFTs were 0.2-1.6.degree.C lower than the lower
     crit. soln. temps. (LCSTs) of the copolymers free in water and the
     differences increased with increasing PVAm content. The catalytic
     activity of the Pt nanoparticles was investigated in the aq.
    hydrogenation of allyl alc. It was found that the
     activity was regulated through temp. - and pH-induced phase sepn. The PVAm
     content also strongly effected the catalytic activity and the morphol. of
     phase sepd. catalysts. With a PVAm content of 4.1 mol%, the
     colloidal platinum sol reversibly changed its catalytic activity with
     changes in temp.
              THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 44
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
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L29 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS AN 2000:566746 CAPLUS

- DN 134:117443
- TI Hydrogenation of olefins in aqueous phase, catalyzed by ligand/protected and polymer-protected rhodium colloids
- AU Borsla, A.; Wilhelm, A. M.; Canselier, J. P.; Delmas, H.
- CS Laboratoire de Genie Chimique-UMR CNRS 5503 (INPT/UPS) Ecole Nationale Superieure d'Ingenieurs de Genie Chimique, Toulouse, 31078/4, Fr.
- SO Studies in Surface Science and Catalysis (2000), 130C(International Congress on Catalysis, 2000, Pt. C), 2093-2098

 CODEN: SSCTDM; ISSN: 0167-2991
- PB Elsevier Science B.V.
- DT Journal
- LA English
- M-tri-sulfonated triphenylphosphine oxide as its sodium salt (OTPPTS) and AB highly water sol. polymers such as poly(vinyl alc.) (PVA) and poly(vinylpyrrolidone) (PVP) were used to stabilize colloidal suspensions of active rhodium particles. The stabilized colloids were used as catalysts in hydrogenation of oct-1-ene in a two-liq. phase system. The effect of various parameters on the stability and activity of the metal nanoparticles under more or less severe conditions was studied. For OTPPTS-protected rhodium colloid systems, the colloid stability improved by increasing the P/Rh molar ratio. Lowering the pressure also improved stability, but led to lower catalytic activity while temp. had no effect on stability but rather on the formation of the active species. The PVP(K15)-Rh colloids can be re-used more than one time at 50.degree. and 0.3 MPa without loss of activity, but not the PVP(K30)-Rh and PVA-Rh colloids. Recycling of the catalytic phase provided unchanged turnover frequencies. Suppression of co-solvent did not result in noticeable changes in activity, indicating that the reaction takes place at the interface. The MET micrographs of the catalytic phase of the OTPPTS-Rh system after use in hydrogenation shows 4 nm particles contg. rhodium oxide and metallic rhodium. IR spectroscopy anal. of the PVP-Rh colloids shows geminal and terminal Rh-CO species with a preponderance of the geminal ones.
- RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L29 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS
- AN 1999:150983 CAPLUS
- DN 130:257756
- TI Homogeneous hydrogenation catalysis with monodisperse, dendrimer-encapsulated Pd and Pt nanoparticles
- AU Zhao, Mingqi; Crooks, Richard M.
- CS Department of Chemistry, Texas A and M University, College Station, TX, 77842-3012, USA
- SO Angewandte Chemie, International Edition (1999), 38(3), 364-366 CODEN: ACIEF5; ISSN: 1433-7851
- PB Wiley-VCH Verlag GmbH
- DT Journal
- LA English
- AB In this report we show that composite materials that consist of noble metal nanoparticles stabilized within

dendrimer interiors are suitable for use as homogeneous

hydrogenation catalysts. These interesting new

materials are prepd. by sorbing PdII or PtII ions into hydroxyl-terminated poly(amidoamine) (PAMAM) dendrimers (Gn-OH, were Gn represents the nth generation) where they complex with interion amine groups. Subsequent chem. redn. of the **metal** ions with BH4- yields

dendrimer-encapsulated metal nanoparticles that

contain the same no. of atoms as were preloaded into the dendrimer initially. The resulting composites are sol. in water and stable, either dry or solvated, for at least several months. Thus, the dendrimer acts as both a template for the prepn. of monodisperse nanoparticles and a porous stabilizer. Dendrimer-encapsulated Pd clusters exhibit

high catalytic activity for the hydrogenation of alkenes in water. Importantly, the catalytic activity can be controlled by adjusting the size (generation) of the dendrimer; i.e., the dendrimer acts as a "nanofilter" with a synthetically controllable mesh. THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 16 ALL CITATIONS AVAILABLE IN THE RE FORMAT => s 127 and carbohydrate 108745 CARBOHYDRATE 116172 CARBOHYDRATES 176447 CARBOHYDRATE (CARBOHYDRATE OR CARBOHYDRATES) 0 L27 AND CARBOHYDRATE L30 => s 127 and sugar 211736 SUGAR 113563 SUGARS 277596 SUGAR (SUGAR OR SUGARS) L31 0 L27 AND SUGAR => s 127 and fructose 53219 FRUCTOSE 82 FRUCTOSES 53232 FRUCTOSE (FRUCTOSE OR FRUCTOSES) 0 L27 AND FRUCTOSE L32 => s 127 and glucose 341326 GLUCOSE 712 GLUCOSES 341480 GLUCOSE (GLUCOSE OR GLUCOSES) L33 0 L27 AND GLUCOSE => s 127 and sorbose 3097 SORBOSE 38 SORBOSES 3108 SORBOSE (SORBOSE OR SORBOSES) 0 L27 AND SORBOSE L34 => s 120 and reduction 262772 REDUCTION 5081 REDUCTIONS 265419 REDUCTION (REDUCTION OR REDUCTIONS) 742417 REDN 39209 REDNS 766916 REDN (REDN OR REDNS) 891539 REDUCTION (REDUCTION OR REDN) L35 15 L20 AND REDUCTION => s 135 and aqueous 148293 AQUEOUS 1 AQUEOUSES

(AQUEOUS OR AQUEOUSES)

148294 AQUEOUS

933909 AQ 119 AQS 933982 AQ L36 3 L35 AND AQUEOUS

=> dis 136 1-3 bib abs

L36 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS

2001:442833 CAPLUS AN

135:182330 DN

- The Effect of Stabilizers on the Catalytic Activity and ΤI Stability of Pd Colloidal Nanoparticles in the Suzuki Reactions in Aqueous Solution
- ΑU Li, Yin; El-Sayed, Mostafa A.
- Laser Dynamics Laboratory School of Chemistry and Biochemistry, Georgia CS Institute of Technology, Atlanta, GA, 30332-0400, USA
- Journal of Physical Chemistry B (2001), 105(37), 8938-8943 SO CODEN: JPCBFK; ISSN: 1089-5647
- PB American Chemical Society
- DT Journal
- LA English
- The role was studied, of capping materials on catalytic activity and AΒ stability of transition ${\tt metal}$ nanoparticles used in catalysis in soln. The Pd nanoparticles were prepd. by redn. of metal salts in the presence of stabilizers, hydroxy-terminated poly(amido-amine) (PAMAM) dendrimers (Gn-OH, where Gn represents the nth generation), polystyrene-b-poly(sodium acrylate), and poly(N-vinyl-2-pyrrolidone) (PVP). The particles were used as catalysts in Suzuki reactions in an aq. medium to study the effects of these stabilizers on the metallic nanoparticle catalytic activity and stability. The stability of the Pd nanoparticles was measured by the tendency of nanoparticles to give Pd black powder after the catalytic reaction. The Suzuki reaction is a good acid test for examg. the stability of these nanoparticles, as it takes place when refluxed at about 100.degree. for 24 h. The stability depends on the type of stabilizer, reactant, and base used in the reaction system. The Pd nanoparticles stabilized with the block copolymer, G3 dendrimer, and PVP are efficient catalysts for the Suzuki reactions between phenylboronic acid (or 2-thiopheneboronic acid) and iodobenzene. The G4 dendrimer is an effective stabilizer; however, strong encapsulation of Pd particles in the dendrimer results in a loss of catalytic activity. The Suzuki reactions between arylboronic acids and bromoarenes catalyzed by Pd nanoparticles result in byproducts due to the homo-coupling of bromoarenes. The two properties are anti-correlated, i.e., the most stable is the least catalytically active.
- THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 42 ALL CITATIONS AVAILABLE IN THE RE FORMAT
- ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS
- ΑN 2001:65241 CAPLUS
- DN 134:268058
- TT Synthesis and functionalities of poly(N-vinylalkyloamide). XIII. Temperature and pH dependence of the catalytic activity of colloidal platinum nanoparticles stabilized by poly[(vinylamine)-co-(N-vinylisobutyramide)]
- ΑU Chen, Chun-Wei; Arai, Kumiko; Yamamoto, Kazuya; Serizawa, Takeshi; Akashi, Mitsuru
- CS Dep. Applied Chem. Chemical Eng., Kagoshima Univ., Kagoshima, 890-0065,
- Macromolecular Chemistry and Physics (2000), 201(18), 2811-2819 SO CODEN: MCHPES; ISSN: 1022-1352
- PΒ Wiley-VCH Verlag GmbH

DT Journal

LA English

AΒ Colloidal platinum nanoparticles in the size range of 5-35 .ANG. have been successfully prepd. in water at room temp. by NaBH4 redn . of ionic platinum in the presence of poly[(vinylamine)-co-(Nvinylisobutyramide)] (PVAm-co-PNVIBA). The temp. - and pH-responsive copolymer was used for the first time as the stabilizer of colloidal metal particles. Three PVAm-co-PNVIBA copolymers with PVAm contents of 4.1, 8.3, and 19.8 mol-% were examd. The particle size and morphol. of the platinum colloids varied with the copolymer compn., as confirmed by TEM measurements. The polymer-stabilized Pt nanoparticles pptd. on heating above their crit. flocculation temps. (CFTs), which were strongly dependent on the soln. pH and the copolymer compn. The CFTs were 0.2-1.6.degree.C lower than the lower crit. soln. temps. (LCSTs) of the copolymers free in water and the differences increased with increasing PVAm content. The catalytic activity of the Pt nanoparticles was investigated in the aq. hydrogenation of allyl alc. It was found that the activity was regulated through temp. - and pH-induced phase sepn. The PVAm content also strongly effected the catalytic activity and the morphol. of phase sepd. catalysts. With a PVAm content of 4.1 mol%, the colloidal platinum sol reversibly changed its catalytic activity with changes in temp.

RE.CNT 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS

AN 1999:760465 CAPLUS

DN 132:107585

TI Preparation and characterization of solvent-stabilized nanoparticulate platinum and palladium and their catalytic behavior towards the enantioselective hydrogenation of ethyl pyruvate

AU Collier, P. J.; Iggo, J. A.; Whyman, R.

CS Department of Chemistry, University of Liverpool, Liverpool, UK SO Journal of Molecular Catalysis A: Chemical (1999), 146(1-2), 149-157 CODEN: JMCCF2; ISSN: 1381-1169

PB Elsevier Science B.V.

DT Journal

LA English

AB

Solvent-stabilized Pt and Pd nanoparticles, of s size range 2.3-2.8 nm and 2.7-3.8 nm, resp., were prepd. by metal vapor synthesis routes, characterized by TEM, and their behavior as catalysts for the enantioselective hydrogenation of Et pyruvate (EP) studied; comparisons were effected with the performance of std. supported Pt and Pd catalysts. Cinchona alkaloid-modified Pt nanoparticles display parallel behavior to that exhibited by their conventional supported counterparts both in terms of the sense of the enantioselectivity in the Et lactate product and in the acceleration in reaction rate relative to the unmodified system. With Pd, however, significant differences are noted. Here, the sense of the enantioselectivity relative to that reported previously over conventional supported catalysts is reversed, i.e., an (R) - vs. (S)-enantiomer switch occurs, and a rate acceleration rather than retardation is noted on cinchona alkaloid modification. The Pt particle size distribution shows a higher degree of monodispersity after use in catalysis, although the av. particle size remains essentially unchanged, whereas the behavior of the Pd nanoparticles shows evidence of concn. dependence, lower concns. showing Pt-like behavior but more highly concd. prepns. showing evidence of significant aggregation during catalysis. With Pt catalysts, the presence of H2O as a component of the ketonic solvent system results in a significant acceleration in overall reaction rate with both conventional supported catalysts and their solvent-stabilized counterparts. In sharp contrast, totally aq.-based colloidal Pt prepns., obtained

by conventional salt **redn**., display very low reaction rates and enantioselectivities.

NT 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT RE.CNT 19

ANSWER 1 OF 5 MEDLINE

IN-PROCESS ACCESSION NUMBER: 2002478532 DOCUMENT NUMBER: 22225843 PubMed ID: 12240156

Nanoparticles as structural and functional units TITLE:

in surface-confined architectures.

AUTHOR: Shipway A N; Willner I

Institute of Chemistry, Hebrew University of Jerusalem, CORPORATE SOURCE:

Jerusalem 91904, Israel.

SOURCE: Chem Commun (Camb), (2001 Oct 21) (20) 2035-45.

Journal code: 9610838. ISSN: 1359-7345.

PUB. COUNTRY:

England: United Kingdom DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE: Entered STN: 20020921

Last Updated on STN: 20020921

The nanoscale engineering of functional chemical assemblies has attracted recent research effort to provide dense information storage, miniaturized sensors, efficient energy conversion, light-harvesting, and mechanical motion. Functional nanoparticles exhibiting unique photonic, electronic and catalytic properties provide invaluable building blocks for such nanoengineered architectures. Metal nanoparticle arrays crosslinked by molecular receptor units on electrodes act as selective sensing interfaces with controlled porosity and tunable sensitivity. Photosensitizer/electron-acceptor bridged arrays of Au-nanoparticles on conductive supports act as photoelectrochemically active electrodes. Semiconductor nanoparticle composites on surfaces act as efficient light collecting systems, and nanoengineered semiconductor 'core-shell' nanocrystal assemblies reveal enhanced photoelectrochemical performance due to effective charge separation. Layered metal and semiconductor nanoparticle arrays crosslinked by nucleic acids find applications in the optical, electronic and photoelectrochemical detection of DNA. Metal and semiconductor nanoparticles assembled on DNA templates may be used to generate complex electronic circuitry. Nanoparticles incorporated in hydrogel matrices yield new composite materials with novel magnetic, optical and electronic properties.

ANSWER 2 OF 5 MEDLINE

ACCESSION NUMBER: 2000194022 MEDLINE

DOCUMENT NUMBER: 20194022 PubMed ID: 10673315

TITLE: Hybrid electrochemical/chemical synthesis of quantum dots.

Penner R M AUTHOR:

CORPORATE SOURCE: Institute for Surface and Interface Science, Department of

Chemistry, University of California, Irvine, California

92679-2025, USA.. rmpenner@uci.edu

Acc Chem Res, (2000 Feb) 33 (2) 78-86. SOURCE:

Journal code: 0157313. ISSN: 0001-4842.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200005

ENTRY DATE: Entered STN: 20000512

> Last Updated on STN: 20000512 Entered Medline: 20000501

AB The "electrochemical/chemical method" (or "E/C method") is a new wet chemical method for synthesizing semiconductor quantum dots on graphite surfaces. The E/C synthesis of quantum dots composed of the generic semiconducting salt, MX, typically involves three steps: (1)

electrochemical deposition of nanoparticles of the metal , M degrees, from a solution of metal ions, M(n)(+); (2) electrochemical oxidation of these metal particles to MO(n)()(/2), and; (3) displacement of the oxygen from MO(n)()(/2) using HX (for example) to yield nanoparticles of MX. This conversion from metal to metal oxide to metal salt occurs on a particle-by-particle basis; that is, each metal nanoparticle is converted into a semiconductor nanoparticle. E/C-synthesized beta-CuI and CdS quantum dots possess many of the attributes of quantum dots synthesized using molecular beam epitaxy, including epitaxial orientation on the graphite surface, a narrow size dispersion, and strong, particle size-tunable photoluminescence. However, the E/C method is faster, cheaper, and applicable to a greater number of materials.

ANSWER 3 OF 5 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2002263725 EMBASE

TITLE: Plasmon-assisted transmission of entangled photons.

AUTHOR: Altewischer E.; Van Exter M.P.; Woerdman J.P.

CORPORATE SOURCE: E. Altewischer, Leiden University, Huygens Laboratory, PO

Box 9504, 2300 RA Leiden, Netherlands.

erwin@molphys.leidenuniv.nl

SOURCE: Nature, (18 Jul 2002) 418/6895 (304-306).

Refs: 16

ISSN: 0028-0836 CODEN: NATUAS

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 027 Biophysics, Bioengineering and Medical

Instrumentation

LANGUAGE: English SUMMARY LANGUAGE: English

The state of a two-particle system is said to be entangled when its quantum-mechanical wavefunction cannot be factorized into two single-particle wavefunctions. This leads to one of the strongest counter-intuitive features of quantum mechanics, namely non-locality. Experimental realization of quantum entanglement is relatively easy for photons; a starting photon can spontaneously split into a pair of entangled photons inside a nonlinear crystal. Here we investigate the effects of nanostructured metal optical elements on the properties of entangled photons. To this end, we place optically thick metal films perforated with a periodic array of subwavelength holes in the paths of the two entangled photons. Such arrays convert photons into surface-plasmon waves-optically excited compressive charge density waves-which tunnel through the holes before reradiating as photons at the far side. We address the question of whether the entanglement survives such a conversion process. Our coincidence counting measurements show that it does, so demonstrating that the surface plasmons have a true quantum nature. Focusing one of the photon beams on its array reduces the quality of the entanglement. The propagation of the surface plasmons makes the array effectively act as a 'which way' detector.

L4 ANSWER 4 OF 5 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2002187476 EMBASE

TITLE: Nanoporous iron oxide membranes: Layer-by-layer deposition

and electrochemical characterisation of processes within

nanopores.

AUTHOR: McKenzie K.J.; Marken F.; Hyde M.; Compton R.G.

CORPORATE SOURCE: F. Marken, Department of Chemistry, Loughborough

University, Leicestershire LE11 3TU, United Kingdom.

f.marken@lboro.ac.uk

SOURCE: New Journal of Chemistry, (2002) 26/5 (625-629).

Refs: 27

ISSN: 1144-0546 CODEN: NJCHE5

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

AB A versatile procedure for the formation of nanoporous **metal** oxide membranes is reported, based on a layer-by-layer deposition

procedure ('directed assembly') of metal oxide nanoparticles with appropriate 'linker' molecules; here Fe(2)O(3) particles and phytic acid. Two types of nanoporous Fe(2)O(3) membranes have been prepared and characterised: (A) a nanofilm deposit composed of 4-5 nm diameter Fe(2)O(3) particles linked by phytic acid and (B) a nanoporous film formed after calcination of the type A deposit at 500.degree.C in air. The nanofilm deposits are characterised by microscopy (SEM and AFM) and by electrochemical methods. Mechanically stable and homogeneous nanofilm deposits with controlled thickness (ca. 3 nm per layer deposited) were obtained. Transport of small molecules or ions through the nanoporous structure and their electrochemical conversion are shown to be fast in the presence of a sufficiently high concentration of supporting electrolyte. During the electrochemical oxidation of ferrocyanide, Fe(CN)(6)(-4), the nanoporous structure of the type A deposit is shown to act as an 'active' membrane, which changes the electrode kinetics by 'double-layer superposition' effects. For the second type of nanofilm, type B, ferrocyanide is accumulated by adsorption within the porous structure.

L4 ANSWER 5 OF 5 EMBASE COPYRIGHT 2002 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2001388918 EMBASE

TITLE: Nanoparticles as structural and functional units

in surface-confined architectures.

AUTHOR: Shipway A.N.; Willner I.

CORPORATE SOURCE: I. Willner, Institute of Chemistry, Hebrew University of

Jerusalem, Jerusalem 91904, Israel. willnea@vms.huji.ac.il

SOURCE: Chemical Communications, (21 Oct 2001) 7/20 (2035-2045).

Refs: 150

ISSN: 1359-7345 CODEN: CHCOFS

COUNTRY: United Kingdom

DOCUMENT TYPE: Journal; General Review

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

properties.

The nanoscale engineering of functional chemical assemblies has attracted recent research effort to provide dense information storage, miniaturized sensors, efficient energy conversion, light-harvesting, and mechanical motion. Functional nanoparticles exhibiting unique photonic, electronic and catalytic properties provide invaluable building blocks for such nanoengineered architectures. Metal nanoparticle arrays crosslinked by molecular receptor units on electrodes act as selective sensing interfaces with controlled porosity and tunable sensitivity. Photosensitizer/electron-acceptor bridged arrays of Au-nanoparticles on conductive supports act as photoelectrochemically active electrodes. Semiconductor nanoparticle composites on surfaces act as efficient light collecting systems, and nanoengineered semiconductor 'core - shell' nanocrystal assemblies reveal enhanced photoelectrochemical performance due to effective charge separation. Layered metal and semiconductor nanoparticle arrays crosslinked by nucleic acids find applications in the optical, electronic and photoelectrochemical detection of DNA. Metal and semiconductor nanoparticles assembled on DNA templates may be used to generate complex electronic circuitry. Nanoparticles incorporated in hydrogel matrices yield new composite materials with novel magnetic, optical and electronic